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54) Title: GAS-PERMEABLE MEMBRANE 87) Abstract		
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GAS-PERMEABLE MEMBRANE

This invention relates to gas-permeable membranes and their use in packaging, especially the packaging of fresh produce.

Fresh cut fruit and vegetables, and other respiring biological materials consume oxygen (O2) and produce carbon dioxide (CO2), at rates which depend upon temperature and the stage of their development. Their storage stability depends on the relative and absolute concentrations of O₂ and CO₂ in the atmosphere surrounding them, and on temperature. Ideally, a respiring material should be stored in a container whose permeability to O2 and CO2 is correlated with (i) the atmosphere outside the package, (ii) the rates at which the material consumes O2 and produces CO2, and (iii) the temperature, to produce an atmosphere within the container having O2 and CO2 concentrations equal to the optimum values for preservation of the material. The permeability to water vapor may also be significant. This is the principle behind the technology of controlled atmosphere packaging (CAP) and modified atmosphere packaging (MAP), as discussed, for example, in US Patent Nos. 4,734,324 (Hill), 4,830,863 (Jones), 4,842,875 (Anderson), 4,879,078 (Antoon), 4,910,032 (Antoon), 4,923,703 (Antoon), 5,045,331 (Antoon), 5,160,768 (Antoon) and 5,254,354 (Stewart), and European Patent Applications Nos. 0,351,115 and 0,351,116 (Courtaulds). The disclosure of each of these publications is incorporated herein by reference.

The preferred packaging atmosphere depends on the stored material. For example, some materials, e.g. broccoli, are best stored in an atmosphere containing 1-2% O_2 and 5-10% CO_2 . For other materials, an atmosphere containing 1-2% O_2 and 12-30% CO_2 , e.g. about 15% CO_2 , is preferred. Thus, CO_2 concentrations of 10 to 30% slow the respiration rate of some fruit and reduce the activity of some decay-causing organisms; for example, a CO_2 concentration of 20% delays grey mold decay in rasberries and extends their shelf life.

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Although much research has been carried out, known packaging techniques have many shortcomings for respiring biological materials. We have discovered, in accordance with this invention, that by forming thin polymeric coatings on microporous films, it is possible to create gas-permeable membranes which have novel and desirable combinations of O₂ permeability, change in O₂ permeability with temperature, and ratio of CO₂ permeability to O₂ permeability. Improved results can be obtained using a wide range of microporous base films and coating polymers. However, a particular advantage of the present invention is that it makes it possible to design packages which are tailored to the requirements of particular respiring materials. As further discussed below, the gasperneable membranes of this invention are generally used as control sections which provide the sole, or at least the principal, pathway for gases to enter or leave a sealed container containing a respiring material.

In describing the invention below, the following abbreviations, definitions, and methods of measurement are used. OTR is O2 permeability. COTR is CO2 permeability. OTR and COTR values are given in ml/m2.atm.24 hrs, with the equivalent in cc/100 inch2.atm.24 hrs given in parentheses. OTR and COTR were measured using a permeability cell (supplied by Millipore) in which a mixture of O2, CO2 and helium is applied to the sample, using a pressure of 0.7 kg/cm2 (10 psi) except where otherwise noted, and the gases passing through the sample were analyzed for O2 and CO3 by a gas chromatograph. The cell could be placed in a water bath to control the temperature. The abbreviation P10 is used to denote the ratio of the oxygen permeability at a first temperature T1°C to the oxygen permeability at a second temperature T2, where T2 is (T1-10)°C, T₁ being 10°C and T₂ being 0°C unless otherwise noted. The abbreviation R is used to denote the ratio of CO2 permeability to O2 permeability, both permeabilities being measured at 20°C unless otherwise noted. Pore sizes given in this specification are measured by mercury porosimetry or an equivalent procedure. Parts and percentages are by weight, temperatures are in degrees Centigrade, and molecular weights are weight average molecular weights expressed in Daltons. For crystalline polymers, the

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abbreviation T_o is used to denote the onset of melting, the abbreviation T_p is used to denote the crystalline melting point, and the abbreviation ΔH is used to denote the heat of fusion. T_o , T_p and ΔH are measured by means of a differential scanning calorimeter (DSC) at a rate of $10^oC/m$ inute and on the second heating cycle.

Typically, a microporous film has an R ratio of about 1, and OTR and COTR values which (i) are very high, (ii) do not change much with the thickness of the film, and (iii) do not change much with temperature (leading to P₁₀ ratios of about 1). A continuous polymeric layer, on the other hand, typically has an R ratio substantially greater than 1 (generally 2 to 6, depending on the polymer itself), and has OTR and COTR values which (i) are relatively low, (ii) are inversely proportional to the thickness of the layer, and (iii) change substantially with temperature (leading to P₁₀ ratios substantially greater than 1, generally at least 1.3). At practical thicknesses, such continuous polymeric layers have OTR and COTR values which are undesirably low.

We have discovered that when a membrane is prepared by coating a thin layer of a polymer onto a suitable microporous film, it has permeability characteristics which depend on both the coating polymer and the microporous film. We do not know exactly why this is so, and the results achieved by this invention do not depend upon any theory of its operation. However, we believe that the coating polymer effectively blocks most, but not all, of the pores of the microporous film (with the smaller pores being preferentially blocked); and that as a result, the permeability of the membrane results in part from gases which pass through the unblocked pores and in part from gases which pass through the coating polymer. In any event, the invention makes it possible to prepare nevel membranes having very desirable permeability characteristics, and to achieve controlled variation of those characteristics. For example, the invention makes it possible to prepare membranes having an OTR greater than 775,000 (50,000), e.g. 1,550,000 (100,000) to 3,875,000 (250,000), or even higher, e.g. up to 7,750,000 (500,000) or more, a P₁₀ ratio of at least 1.3, e.g. at least 2.6, and an R ratio of at least 1.5, e.g. at least 2.0.

The microporous film and the coating polymer must be selected and correlated to produce a membrane having particular properties, but those skilled in the art will have no difficulty, having regard to the disclosure in this specification and their own knowledge, in achieving a wide range of useful results.

The size and distribution of the pores in the microporous film are important factors. If the pores are too small, the coating polymer tends to form a continuous layer which is either too thin to be durable under routine handling, or too thick to have an acceptable OTR. If the pores are too large, the coating polymer may be unable to bridge over them, so that the coating polymer plays little or no part in determining the permeability characteristics of the membrane. This may happen even if the average pore size is relatively low, if the pores have a wide range of sizes; for example the coating polymer may effectively block many of the pores, but still fail to block the larger pores, whose permeability then dominates the permeability of the membrane as a whole.

The roughness of the microporous film can also be an important factor. The coating weight of the coating polymer must be very small, and in consequence the thickness of the coating polymer is also very small. If such a thin layer is in intimate contact with an irregular surface, it is more likely to be able to withstand abrasive forces during use than a layer of the same thickness which lies on a relatively smooth surface.

The coating polymer should be selected so that the membrane has a desired P_{10} ratio and a desired R ratio, and should be coated onto the microporous film at a coating weight which results in a membrane having the desired balance between the permeability characteristics of the microporous film and of the coating polymer. For example, by choosing a crystalline coating polymer whose T_p is within or a little below an expected range of storage temperatures, it is possible to produce a membrane whose P_{10} is relatively large in the storage temperature range; furthermore, the size of the P_{10} ratio can be increased by increasing the AH of the coating polymer. Similarly, a membrane having

a relatively large (or small) R ratio can be produced by selecting a coating polymer having a relatively high (or small) inherent R ratio. In this way, the invention makes it possible to produce membranes whose properties can be tailored, much more closely than is now possible, to the needs of a particular respiring biological material.

In a first preferred aspect, this invention provides a gas-permeable membrane which comprises

- (a) a microporous polymeric film, and
- a polymeric coating on the microporous film, the polymeric coating changing the permeability of the microporous film so that the membrane
 - has a P₁₀ ratio, over at least one 10°C range between -5 and 15°C, of at least 1.3:
 - has an oxygen permeability (OTR), at all temperatures between 20° and 25°C, of at least 775,000 ml/m².atm.24 hrs (50,000 cc/100 inch².atm.24 hrs); and
 - (iii) has an R ratio of at least 1.5;

the P10, OTR and R values being measured at a pressure of 0.035 kg/cm2.

 $\label{eq:incomposition} In a second preferred aspect, this invention provides a package which is stored in $20 air and which comprises $20 are and which comprises $20 are all 20 are a$

- (a) a sealed container, and
- within the sealed container, a respiring biological material and a packaging atmosphere around the biological material

the sealed container including one or more permeable control sections which provide substantially the only pathways for oxygen and carbon dioxide to enter or leave the packaging atmosphere, at least one said permeable control section being a gas-permeable membrane as defined in the first aspect of the invention.

Microporous Base Films

The microporous base film used in this invention comprises a polymeric matrix which provides a network of interconnected pores such that gases can pass through the film. The average pore size of the base film can be 0.02 to 5 micron, but is preferably greater than 0.05 micron and less than 0.24, particularly less than 0.20, especially less than 0.15, micron. Preferably at least 70%, particularly at least 90%, of the pores have a pore size of less than 0.24 micron. Preferably, at least 60%, particularly at least 80%, of the pores have a a pore size of less than about 0.15 micron. Particularly at least 60%, especially at least 70%, of the pores have a pore size of less than about 0.11 micron. The pores preferably constitute 35 to 80%, particularly 60 to 75%, by volume of the film. A narrow range of pore sizes is preferred. For example, it is preferred that less than 20% of the pores have a size less than 0.014 micron, and that less than 20% of the pores have a pore size greater than 0.13 micron.

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It is highly desirable that the base film should have sufficient strength (and other physical properties) to ensure that it is not damaged in processing or in use as part of a container. Accordingly, the base film preferably has a tear strength of at least 30g, particularly at least 70g, when measured by the following test. A sample of the film, 10 by 1.27 cm, is used. A slit 5 mm long is made in one of the short sides. The two flaps on either side of the slit are placed in opposing jaws of an Instron tensile tester, and the tear strength measured at a jaw separation rate of 12.7 cm (5 inch)/minute. If the base film has too low a tear strength, it is possible to prepare a satisfactory membrane by laminating the base film, before or after it has been coated, to a support film which has a satisfactory tear strength and sufficient porosity that its presence does not adversely affect the permeability of the coated membrane. However, the added cost of this expedient is highly undesirable.

It is also preferred that the microporous base film should have a Sheffield smoothness of at least 30. It is also preferred that the base film is composed of a material that can be heat-scaled.

Preferred polymers for polymeric matrix of the base film are

- essentially linear ultrahigh molecular weight polyethylene having an intrinsic viscosity of at least 18, preferably 18 to 39, deciliters/gram,
- essentially linear ultrahigh molecular weight polypropylene having an intrinsic viscosity of at least 6 deciliters/gram, and
- (3) mixtures of (1) and (2).

The preferred base film includes a finely divided, particulate, substantially waterinsoluble, inorganic filler, for example a siliceous filler, which is distributed throughout
the matrix and which is present in amount 50 to 90%, particularly 50 to 85%, by weight
of the base film. The filler is preferably silica, particularly precipitated silica, especially
silica having an average ultimate particle size of less than 0.1 micrometer. Preferably the
filler occupies 35 to 80% of the total volume of microporous film. Particularly preferred,
because they have a relatively narrow range of pore sizes, are films made by stretching a
film which contains an inorganic filler and a processing oil, e.g. a parafinic oil,
naphthenic oil or aromatic oil, uniformly distributed therein. Suitable films are disclosed,
for example, in US Patent Nos. 4,937,115 and 3,351,495, the disclosures of which are
incorporated herein by reference. Such films are sold by PPG Industries under the
tradename "Teslin".

The thickness of the base film is preferably 0.03 to 0.65 mm.

The microporous base film can be modified by calendering at a nip pressure of 17.8 to 266.6 kg per linear cm (100 to 1500 pli). We have found that calendering the base film results in coated films which have reduced OTR values and increased R values. The base film can be uniaxially or biaxially calendered. The base film can also be stretched, uniaxially or biaxially.

Coating Polymers

The coating polymer can be a single polymer or a mixture of two or more different polymers. Preferably the coating polymer is a crystalline polymer having a Tn of -5 to +40°C, particularly -5 to 15°C, particularly 0 to 15°C, e.g. 1° to 15°C, and a ΔH of at least 5 J/g, particularly at least 20 J/g. We have found that the higher the ΔH of the polymer, the higher its P10 value over temperature ranges which include T2. The T2 and T2 values of the polmer are preferably such that T_n-T_n is less than 10°C, particularly 5 to 10°C. Suitable polymers include those described in US Patent No. 5,254,354, the disclosure of which is incorporated herein by reference. Particularly preferred are side chain crystalline (SCC) polymers. SCC polymers can be prepared for example by copolymerizing (i) at least one n-alkyl acrylate or methacrylate in which the n-alkyl group contains at least 12 carbon atoms, preferably one or more of docosanyl, octadecyl, hexadecyl, tetradecyl, and dodecyl acrylates, and (ii) one or more comonomers selected from acrylic acid, methacrylic acid, and esters of acrylic or methacrylic acid in which the esterifying group contains less than 10 carbon atoms, e.g. hydroxyethyl butyl, hexyl, or 2-ethyl hexyl. These SCC polymers generally have a high R ratio, e.g. greater than 5. Other polymers which can be used include cis-polybutadiene, poly(4-methylpentene), polydimethyl siloxanes, and ethylene-propylene rubbers. The polymer can be crosslinked after it has been applied, for example to reduce its tackiness.

Coating Compositions

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The coating polymer is preferably applied to the porous substrate as a coating composition which comprises the coating polymer dissolved in a suitable solvent, e.g. tohene, tetrahydrofuran, heptane or methyl ethyl ketone. After the composition has been applied, the solvent is removed by heating, leaving the polymer adherent to the substrate. The concentration of the coating polymer in the coeting composition is preferably 2 to 12%, e.g. 5 to 10%, based on the weight of the composition. The coating composition can contain other ingredients in addition to the polymer and the solvent, for example a

crosslinking agent which is activated after the solvent has been removed, e.g. aluminum acetyl acetonate or a polyfunctional aziridine such as the product available from Aldrich Chemicals under the tradename XAMA 7.

Application of the Coating Compositions

The coating can be carried out in any convenient way, for example by hand, using a Meyer rod, or using commercially available coating equipment, e.g. gravure-coating equipment, which is preferred, or knife-over-roll equipment. A single coating is normally adequate, but a second coating of the same or a different coating composition can be applied after drying the first. Preferably the coating is carried out using a gravure coating roll having a theoretical cell volume of 31 x 10⁶ to 23.2.5 x 10⁶ cubic microns per mm² (20 x 10⁹ to 150 x 10⁹ cubic microns per inch²), preferably 62 x 10⁶ to 124 x 10⁶ cubic microns per mm² (40 x 10⁹ to 80 x 10⁹ cubic microns per inch²), or using equipment which mrovides a similar coating weight. The coating weight is preferably 1.7 to 2.9 g/m².

Properties of the Membranes

As discussed above, the properties of the membranes of the invention depend upon a number of factors, including the base film, the coating polymer, the coating composition, and the amount of coating composition applied to the base film. The membrane preferably has a P₁₀ ratio, over at least one 10°C range between -5 and 15°C, preferably over at least one 10°C range between 0°C and 15°C, of at least 1.3, preferably at least 2.9, particularly at least 2.5, especially at least 2.6. The membrane preferably has an OTR of 775,000 to 7,750,000 (50,000 to 500,000), preferably 1,550,000 to 3,875,000 (100,000 to 25,000), particularly at least 2,325,000 (150,000). When the OTR is 775,000 to 3,100,000 (50,000 to 200,000), the R ratio of the membrane is preferably at least 2, particularly at least 2.5, especially at least 3; and when the OTR is 775,000 to 3,100,000 (50,000 to 200,000), the R ratio is preferably more than 3.8 - 0.00000045 P' (3.8 - 0.000007P), particularly up to 7.4 - 0.00000116 P' (7.4-0.000018P), especially up to 5.6 -

0.0000084 P (5.6-0.000013P), where P is the OTR in ml/m₂.atm.24 hrs, and P is the OTR in cc/100 inch₂.atm 24 hrs. The membrane preferably also has these values for OTR and R when the OTR and COTR are measured at any temperature between 20°C and 25°C. The membrane preferably has these values for OTR, P₁₀ and R both when the O₂ and CO₂ permeabilities are measured at 0.7 kg/cm² (10 psi) and when they are measured at 0.035 kg/cm² (0.5 psi).

The membrane can be covered by a covering layer, so that the polymeric coating is sandwiched between the microporous film and the covering layer, providing that the covering layer is sufficiently porous not to have an adverse effect on permeability. The covering layer is generally different from, but can be the same as, the base film. The cover layer can be melt-bonded to the coated film. However, the use of a covering layer adds to the cost of the product.

The properties of the membrane can be modified by calendering, uniaxially or biaxially, preferably at a nip pressure of 17.8 to 266.6 kg per linear cm (100 to 1500 pli). We have found that calendering reduces the OTR and increases the R ratio of the membrane.

20 Containers

The properties of the membranes are such that they can be used to control the atmosphere within a container whose walls are relatively impervious to gases except over one or more control sections, the control section, or at least one of the control sections if there are two or more, being provided by a membrane of the invention. In one embodiment, the control section is an aperture which lies in a gas-impervious wall of the container and which is covered by a membrane of the invention. The aperture can be, for example, 5 to 50% of the total wall area of the container. The container can be, for example, a bag of flexible polymeric material, e.g. a laminate of two different polymeric films. Preferably the uncoated side of the membrane is secured, e.g. heat-sealed, impulse-

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scaled, or radio frequency scaled, to the container, around the periphery of the aperture. For this purpose, the outer surface of the container is preferably composed of a material which can be heat-scaled to the base film. In a typical procedure, the membrane is heatscaled to the enclosure using an impulse scaler at 115°C for 0.4 secs.

In another embodiment, the membrane is in the form of a strip which runs down the length of the container, or which otherwise extends across a complete dimension of the container. This has the advantage that the strip can be incorporated into the sheet material from which the bag or other container is made, so that it is not necessary to add individual patches to each container. Another advantage is that the strip is less likely to interfere with graphical design on the container.

The size of the containers can vary considerably. In one category, the container contains up to 2.26 kg (5 lb) of produce. In another category, the containers are much larger, e.g. containing up to 1500 lb (680 kg) of produce.

A wide range of respiring biological materials can be packaged in containers of the invention, including broccoli, cut lettuce, cauliflower, mushrooms, asparagus, and strawherries.

EXAMPLES

The invention is illustrated in the following Examples, a number of which are comparative Examples. In the Examples, the coating polymers used are often referred to by the following abbreviations.

SCC_1-15 and ACP. These acrylate polymers were prepared by polymerizing the monomers and parts by weight thereof shown in Table 1. In Table 1, the following abbreviations are used for the monomers. AA is acrylic acid, MAA is methacrylic acid, EHA is 2-ethylicxyl acrylate, C4A is butyl acrylate, C6A is hexyl acrylate, C6DA is hexyldiacrylate. C12A is dodecyl acrylate. C12A is dodecyl diacrylate. C14A is

tetradecyl acrylate, C16A is hexadecyl acrylate, and C22A is behenyl acrylate. The monomers were copolymerized in a suitable solvent, e.g. toluene or a mixture of heptane and butyl acetate or ethyl acetate, using a suitable initiator, e.g. azobisisobutyronitrile (AIBN). For example, SCC11 was made by mixing the C6A and C14A with 0.31 parts of AIBN in a 4.1 mixture of heptane and ethyl acetate, and maintaining the mixture at 50°C for 2 days, to give a product having a weight average molecular weight of about 970,000 and a T_p of about 12.5°C. SCC 5 was made by polymerizing the C22A alone, functionalizing the resulting homopolymer with isocyanatoethylmethyl methacrylate, and then polymerizing the other monomers with the functionalized homopolymer. The AH values of the SCC12, SCC13, and SCC14 polymers were 0, 35 and 55 J/g respectively.

Table 1

	AA	MAA	EHA	C4A	C6A	C6DA	C12A	C12DA	C14A	C16A	C22A
SCC1	5	-	 -	†-	-	-	47.5	-	-	47.5	-
SCC2	5	-	-	† -	·		65	-	-	30	-
SCC3	-	5	-	-	-	-	47.5		47.5	1-	-
SCC4	5	1.	-	1-	-	-	60	-	•	35	-
SCC5	3	1-	-	-	-	-	20	-	60	-	20
SCC6	-	-	-	-	-	-	65	0.25	•	35	-
SCC7	1	1-	-	-	-	•	64	0.1	•	35	-
SCC8	-	-	<u> -</u>	-	-	-	-	-	100	-	-
SCC9	-	5	-	1-	-	-	47.5	-	47.5	-	-
SCC10	3	-	-	1-	-	-	-	-	57	40	-
SCC11	-	1-	-	•	15	-	-	-	85	-	-
SCC12	-	5	-	<u>†-</u> -	15	-	-	-	80	-	
SCC13	3	† -	† -	-	42	-	-	-	55	-	-
SCC14	-	-	•	-	15	-	-	-	85	-	-
SCC15	 	-	-	1-	-	-	-	-	100	27	-
ACP	6	-	47	47	1-		-	-	-	1-	1-

<u>CisPB</u> is cis-potybutadiene available from Aldrich Chemical Co. <u>Tflex</u> is an aliphatic

polyurethane available under the tradename Tecoflex SG80A from Thermedics Corp. <u>Sil</u>

is a moisture-curable, RTV (room-temperature vulcanizable) polydimethyl siloxane available under the tradename Silastic 734 from Dow Corning Corporation. <u>Kton</u> is a styrene-ethylene-butylene block copolymer available under the tradename Kraton from Shell Chemical Co. <u>TPX</u> is poly-(4-methyl pentene) available under the tradename TPX from Aldrich Chemicals.

These polymers were used in coating solutions of various concentrations. The SCC and ACP polymers were prepared in solution and were diluted to the desired concentration by addition of a suitable solvent, e.g. heptane. The CisPB, the mixtures of CisPB and SCC8, and the Sil polymers were dissolved in toluene; the Tflex polymers were dissolved in tetahydrofuran; and the Kraton polymers were dissolved in toluene.

The substrates used in the Examples are often referred to by the following abbreviations. \underline{CG} is a porous polyethylene film having a porosity of about 55% and a pore size of about 0.25 microns, available under the tradename Celgard K878 from Hoechst Celanese Corp. \underline{MSX} is a porous polyethylene film available under the tradename MSX 1137P from 3M Co. $\underline{Van L}$ is a filled porous polyethylene film available under the tradename Van Leer 10X from Van Leer Corp. $\underline{Teslin.SP7}$ is a filled porous polyethylene films containing about 60% silica, having a thickness of about 0.18 mm (0.007 inch), a tear strength measured as described above of about 90g, a porosity of about 65%, an average pore size of about 0.1 micron and a largest pore size of 4-10 microns. Teslin X457 is similar to Teslin SP7 but is more porous. Both Teslins are available under the tradename Teslin from PPG Industries. When testde by mercury porosimetry, the distribution of pore sizes in Teslin SP7 was as follows.

Pore Size (microns)	.013	.016	.026	.044	.058	.08	.11	.15	.24	.36	.6
% of pores larger than pore size	90%	80%	70%	60%	50%	40%	30%	20%	10%	5%	2%

respectively.

AKZ0 is a porous polypropylene film in which the pores have a size of 0.1 to 0.29 micron, available under the tradename Akzo 1EPP from Enka AG. Delnet is a perforated high density polyethylene film, 0.11 mm (4.5 mil) thick, with an open area of about 36%, available under the tradename Delnet from Applied Extrusion Technologies. BF 915, Roplast, LB710 and CVP are commercially available laminate packing films in which one layer is polyethylene, available from Barrier, Roplast, Golden Eagle and CVP

The procedure used to coat the substrates was to apply the coating solution with a #30 Meyer rod, and to dry the coating in ambient air for 15 minutes and then at 83°C for 30 minutes

Examples 1-10 and C1-4

Examples 1-10 and C1-4 are summarized in Tables 2-4 below. In each of the tests, the substrate identified in Table 2 was coated with a solution containing the identified percentage of the identified coating polymer. The coated substrate was dried, and its permeability to O₂ and CO₂ was measured at the temperature indicated in Table 2. The results are reported in Table 2. In some cases, the P₁₀ ratios were calculated, as shown in Tables 3 and 4. The P₁₀ ratios in Examples 8-10 demonstrate how the increasing AH values of SCC13 - SC15 result in increasing P₁₀ ratios.

The OTR and R values of some of the Examples are shown graphically in Figure 1 of the accompanying drawings. In Figure 1, curve 1 represents comparative Examples 1K to 1N (SCC 1 on Van Leer), curve 2 represents comparative Examples 1H-J (SCC1 on MSX), curve 3 represents Examples 2O-Q (TPX on Teslin SP7), curve 4 represents Examples 1P, Q and R (SCC1 on Teslin SP7), and curve 5 represents Examples 2H to 2K (cis PB + SCC8 on Teslin SP7).

Table 2

Ex No.	Substrate	Coat Poly		Perr	neability	
		Туре	%	OTR	Ratio CO ₂ /O ₂	°C
*1A	CG	SCC1	2	**	1.0	20
*B		1	5	* **	1.0	20
*C			10	**	1.0	20
*D			20	**	1.0	20
*E	MSX	SCC1	2	8.0	1.0	20
*F			5	**	1.0	20
*G			10	**	1.0	20
*H			20	487,475 (31,450)	1.86	20
*I			25	63,689 (4,109)	5.75	20
*J		1	30	44,175 (2,850)	6.5	20
*K	Van L	SCC1	2	403,077 (26,005)	1.32	20
*L		1	5	29,636 (1,912)	3.92	20
*M			10	62,558 (4,036)	2.39	20
*N		1	20	62,310 (4,020)	2.42	20
0	Teslin SP7	SCC1	2	2,413,458 (155,707)	2.95	20
P			5	1,021,140 (65,880)	4.86	20
Q			10	602,547 (38,874)	5,22	20
R			20	425,025 (27,421)	5.48	20
2A	Teslin SP7	Tflex	5	1,076,537 (69,454)	2.34	20
В			7	238,024 (15,368)	5.9	20
C			15	108,856 (7,023)	6.19	20
D	Teslin SP7	Sil	35	1,607,799 (103,729)	2.98	20
E			40		4.13	20
F			50	809,627 (52,234)	4.7	20
Ğ			100	391,824 (25,279)	5.3	20
Н	Teslin SP7	CisPB	2	3,712,994 (239,548)	2.8	20
I	I	1	5	2,451,728 (158,176)	4.39	20
J		T	10	888,987 (57,354)	6.45	20
K		1	20	392,925 (25,350)	7.2	20
L	Teslin SP7	Kton	5	293,570 (18,940)	4.4	20
M			10	116,606 (7,523)	4.7	20
N			20	46,887 (3,025)	5.0	20
0	Teslin SP7	TPX	5	974,841 (62,893)	3.35	20
P	-		5	923,598 (59,587)	3.3	20
Q		1	2	2,798,478 (180,547	2.38	20

^{*}comparative example
**permeability too high to measure

Table 2 (continued)

		Coatin Polym			eability	
Ex. No.	Substrate	Туре	%	OTR	Ratio CO ₂ /O ₂	•℃
3A	Teslin SP7	SCC2	10	637,592 (41,135)	4.94	20
В		SCC3	5	927,861 (59,862)	4.81	20
C		ACP	5	403,728 (26,047)	5.8	20
D		CisPB	5	2,092,934 (135,028)	4.8	20
E		*CisPB + SCC8	5	1,205,559 (77,778)	5.3	20
F		SCC4	2	1,883,389 (121,509)	3.3	20
G	 	SCC4	10	687,688 (44,367)	5.2	20
H		SCC5	-	930,854 (60,055)	4.72	20
I		SCC6	10	812,959 (52,449)	5.10	20
J		SCC7	25	410,440 (26,480)	5.23	20
4A	Teslin SP7	CisPB	5	125,218 (80,786)	6.50	0
В		CisPB	5	1,367,875 (88,250)	6.09	5
C	1	CisPB	5	1,579,000 (101,871)	5.51	10
D		CisPB	5	1,839,323 (118,666)	5.43	15
Е		CisPB	5	2,092,934 (135,028)	4.82	20
F		CisPB	5	2,328,952 (150,255)	4.55	25
G		*CisPB + SCC8	5	320,617 (20,685)	5.5	0
Н		*CisPB + SCC8	5	394,304 (25,439)	5.5	5
I		*CisPB + SCC8	5	512,848 (33,087)	5.4	10
J		*CisPB + SCC8	5	722,455 (46,610)	5.3	15
K		*CisPB + SCC8	5	1,205,559 (77,778)	5.3	20
L		*CisPB + SCC8	5	1,520,379 (98,089)	5,2	25
М		SCC10	5	1,020,628 (65,847)	5.93	22.2

^{*}a mixture of CisPB (80 parts) and SCC8 (20 parts)

Table 2 (continued)

		Coat Polyi		Perm	eability	
Ex. No.	Substrate	Туре	%	OTR	Ratio CO ₂ /O ₂	°C
5A	Teslin SP7	SCC11	6.5	† 386,461 (24,933)	6.06	0
В			6.5	†521,141 (33,622)	6.16	5
C			6.5	†1,048,590 (67,651)	6.12	10
D			6.5	†1,474,933 (95,157)	5.78	15
Е			6.5	†1,825,001 (117,742)	5.53	. 20
F		117	6.5	†2,225,753 (143,597)	5.21	25
G			6.5	†2,587,33 (166,926)	4.88	30
6	Teslin SP7	SCC12	6.5	†1,286,500 (83,000)	5.29	22
7A	Teslin X457	SCC13	2	†4,022,699 (259,529)	3.92	22
В			1.5	18,789,414 (567,059)	2.62	22
С			1.3	†13160,662 (849,075)	1.68	22
7D	Teslin SP7	SCC13	1	†8,693,239 (560,854)	2,75	22
Е			1.5	†3,262,486 (210,483)	4.88	22
F			2	†2,766,083 (178,547(5.48	22
G			3	†1,869,005 (120,581)	6.03	22
8	Teslin SP7	SCC13	3	†1,518,643 (97,977)	6.32	22
9		SCC15	6.5	†1,758,196 (113,432)	5.58	22
10		SCC16	3.8	†1,113,241 (71,822)	7.06	22
C1	Roplast	None	-	†3,859 (249)		22
C2	BF915	None	-	†4,836 (312)		22
C3	LB710	None	-	†5,161 (333)		22
C4	CVP	None	-	†5,130 (331)		22

 $[\]uparrow$ OTR and COTR were measured at a pressure of 0.035 kg/cm² (0.5 psi) in Examples 5-10 and C1-C4.

Table 3

Ex No	3A	3B	3C	3D	3E	3F	3G	3H	31	3J
P ₁₀ (0-10°C)	2.79	2.00	1.31	1.26	1.60	2.41	3.13	-	4.72	4.33
P ₁₀ (20-30°C)	١.		-	1.34	2.35	-	-	2.73	-	-)

Table 4

Ex No	4A	G	5A	8	9	10	11	C1	C2	C3	C4
P ₁₀ (0-10°C)	1.26	1.6	†2.71	†1.4	†2.4	†2.8	†3.5	†1.54	†1.82	†1.74	†1.79
P ₁₀ (10-20°C)	1.33	2.35	-	-	-	•	-	-	٠	-	-

 \uparrow OTR's in Examples 5A, 8-10 and C1-C4 were measured at a pressure of 0.035 kg/cm² (0.5 psi).

Example 11

A membrane was produced by coating SCC 10 at 8% concentration onto Testin SP7. The resulting product was tacky to the touch. Second and third membranes were produced in the same way except that a crosslinking agent was added to the coating solution. The crosslinking agent was aluminum acetylacetonate (5%, based on polymer) or a polyfunctional aziridene (5%, based on polymer, available under the trade name XAMA 7 from Virginia Chemicals). The resulting products were much less tacky.

5 Example 12

This example illustrates the effect of calendering the coated microporous film.

The OTR and R values and the thicknesses of a coated microporous film made by coating SCC1 onto Testin SP7 were measured before calendering, after uniaxial calendering at 142 kg/linear cm (800 pli) or after uniaxial calendering at 269 kg/linear cm (1500 pli).

The results are shown in Table 5 below.

Table 5

Nip Pressure kg/lin. cm (pli)	Thickness mm (mils)	OTR	R
0	0.17 (6.9)	1,973,010 (127,291)	3.82
142 (800)	0.16 (6.3)	1,742,138 (112396)	4.09
169 (1500)	0.157 (6.2)	1,485,473.5 (95837)	4.64

Example 13

Example 13 is summarized in Table 6. Fresh broccoli was sealed into storage bags (23 x 28 cm) and stored for 13 days at 45°C, or for 9 days at 45°C and 4 days at 55°C. Four control bags (C5 - C8) were composed entirely of polyethylene. The other four bags (13A - 13D) were bags of the invention and were identical to the control bags except that a hole of 4.76 cm diameter was cut in the polyethylene and sealed by means of a membrane which had an O₂ permeability of 1,224,500 (79,000) and which had been prepared by coating Teslin SP7 with the SCC10 polymer. At the end of the storage period, the O₂ and CO₂ concentrations in the bag were measured. The results are shown in Table 5. The broccoli was then cooked for 1 minute in a mirrowave oven at the "high" setting. The broccoli from all the control bags had a strong and offensive odor and flavor before and after cooking. The broccoli from all the bags of the invention had no unpleasant odor or flavor before or after cooking; the broccoli stored at 45°C for all 13 days was in even better condition than that stored at 45°C and 55°C.

Table 6

	Stor	age	After	Storage
	Days at 45°C	Days at 55°C	% CO ₂	% O ₂
C5	13	0	8.3	0.29
C6	13	0	8.0	0.79
C7	9	4	9.1	0.46
C8	9	4	8.5	0.32
13A	13	0	4.1	2.89
13B	13	0	4.8	10.9
13C	9	4	4.1	1.04
13D	9	4	4.3	1.00

Example 14

The following samples were used in this example.

Sample 14A Teslin SP7 which has been coated with a 10% solution of SCC4.

Sample 14B. A laminate of Sample 14A and Teslin SP7 laminated to the coated face of

Sample 6A at about 50°C, applying pressure by means of a 8.85 kg (4 lb) roller.

Sample 14C Teslin SP7 which has been coated with a 1% solution of SCC13.

<u>Sample 14D</u> A laminate of Sample 14A and AKZO laminated to the coated face of Sample 14A in the same way as for Sample 14B.

The OTR's of samples 14A-D and 7B were measured (1) initially and (2) after folding four times (i.e. to 0.125 of original size) and unfolding. The results are shown in Table 7.

Table 7

OTR	14A	14B	14C	14D	7B
Initial	687,735 (44,370)	871,286 (56,212)	†8,693,237 (560,854)	1,277,711 (82,433)	†8,789,414 (567,059)
After folding	**	852,500 (55,000)	†10,081,122 (650,395)	1,282,857 (82,765)	†9,872,291 (636,922)

^{**} too permeable to measure

Example 15

A sample of AKZO film was heat-welded to Sample 14C. The heat weld was produced by an impulse sealer available from Packaging Aids Inc. under the trade name Sealmaster 420. The force needed to break the resulting bond was more than the tear strength of the AKZO film.

Example 16

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The following samples were used in this Example.

Sample 16A Teslin SP7 which had been coated with a 5% solution of SCC4.

<u>Sample 16B</u> A laminate of Sample 16A and Teslin SP7 in the Teslin being laminated to the coated face of Sample 16A under heat and pressure.

Sample 16C A laminate of Sample 16A and Delnet, the Delnet being laminated to the coated face of Sample 16A under heat and pressure.

[†] OTR's were measured at 0.035 kg/cm² (0.5 psi) in Examples 14C and 7B

The O_2 permeabilities of samples 16B and 16C were measured initially, after folding as in Example 14, and after immersion in tap water for 10 minutes, followed by drying with a paper towel. The results are shown in Table 8.

Table 8

	OTR		
	Initial	After Folding	After Immersion
16B	†871,286 (56,212)	1852,500 (55,000)	†479,213 (30,917)
16C	†522,319 (33,698)	†489,009 (31,549)	†489,009 (31,549)

 $[\]uparrow$ OTR's in Examples 16B and 16C were measured at 0.035 kg/cm² (0.5 psi)

CLATMS

- A gas-permeable membrane which comprises
 - (a) a microporous polymeric film, and
 - a polymeric coating on the microporous film, the polymeric coating changing the permeability of the microporous film so that the membrane
 - has a P₁₀ ratio, over at least one 10°C range between -5 and 15°C, of at least 1.3:
 - (ii) has an oxygon permeability (OTR), at all temperatures between 20° and 25°C, of at least 775,000 ml/m².atm.24 hrs (50,000 cc/100 inch².atm.24 hrs; and
 - (iii) has a CO₂/O₂ permeability ratio(R) of at least 1.5;
 the P₁₀, OTR and R values being measured at a pressure of 0.035 kg/cm² (0.5 psi)
- A membrane according to claim 1 which has at least one of the following characteristics
 - (1) the microporous film has an average pore size of less than 0.24 micron;
 - (2) the microporous film has a tear strength of at least 30g;
 - (3) the microporous film has a Sheffield Smoothness of at least 30;
 - (4) the microporous film comprises a polymeric matrix comprising an essentially linear ultrahigh molecular weight polyethylene having an intrinisc viscosity of at least 18 deciliters/g;
 - (5) the microporous film comprises a polymeric matrix comprising an essentially linear ultrahigh molecular weight polypropylene having an intrinsic viscosity of at least 6 deciliters/a:
 - the microporous film comprises a finely divided, particulate, substantially insoluble filler which is distributed throughout the film;
 - (7) the coating weight of the coating polymer is 1.7 to 2.9 g/m²; and
 - (8) the coating polymer is a crystalline polymer having a T_p of -5° to 15°C.

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- A membrane according to claim 1 or 2 which has a P₁₀ ratio between 0 and 10°C of at least 2.6, the P₁₀ being measured at a pressure of 0.035 kg/cm² (0.5 psi).
- 4. A membrane according to claim 1 or 2 which has a P₁₀ ratio, over at least one s 10°C range between -5°C and 15°C, of at least 2.6, an OTR at all temperatures bewteen 20°C and 25°C of at least 1,550,000 ml/m² atm.24 hrs (100,000 cc/inch² atm.24 hrs, and an R ratio of at least 2.5, the P₁₀, OTR and R values being measured at a pressure of 0.035 kg/cm² (0.5 psi).
- 5. A membrane according to claim 1 wherein at least 90% of the pores have a pore size less than 0.24 micron, and which has a P₁₀ ratio, over at least one 10°C range between -5 and 15°C of at least 1.3, an OTR of 775,000 to 3,100,000 ml/m² atm.24 hrs (50,000 to 200,000 cc/100 inch² atm 24 hrs), and an R ratio of at least 2, the P₁₀, OTR and R values being measured at a pressure of 0.7 kg/cm² (10 psi).
 - A membrane according to claim 5 which has at least one of the following characteristics
 - it has a P₁₀ ratio of at least 1.3 over at least one 10°C temperature range between 0° and 15°C;
 - it has a P₁₀ ratio of at least 2 over at least one 10°C temperature range between 0° and 15°C;
 - (3) it has a P₁₀ ratio of at least 2.5 over at least one 10°C temperature range hetween 0° and 15°C;
 - (4) it has an OTR of at least 1,550,000 ml/m².atm.24 hrs (100,000 cc/100 inch².atm.24 hrs);
 - (5) it has an OTR of at least 2,325,000 ml/m².atm.24 hrs (150,000 cc/100 inch² atm.24 hrs):
 - (6) it has an OTR of 775,000 to 3,100,000 ml/m².atm.24 hrs (50,000 to 200,000 cc/100 inch².atm.24 hrs) and an R value of more than (3.8 0,00000045P), where P is the OTR in ml/m².atm.24 hrs;

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- (7) it has an OTR of 775,000 to 3,100,000 ml/m².atm.24 hrs (50,000 to 200,000 cc/100 inch².atm.24 hrs) and an R value of (3.8 0.0000045P) to (7.4 0.000016P), where P is the OTR in ml/m².atm.24 hrs:
- (8) it has an OTR of 775,000 to 3,100,000 ml/m².atm.24 hrs (50,000 to 200,000 cc/100 inch².atm.24 hrs) and an R value of (3.8 0.0000045P) to (5.6 0.000084Pl), where PI is the OTR in ml/m².atm.24 hrs;
- (9) it has an R ratio of at least 2.5; and
- (10) it has an R ratio of at least 3.

the P_{10} , OTR and R values being measured at a pressure of 0.07 kg/cm² (10 psi).

- 7. A membrane according to claim 5 or 6 wherein the microporous film has at least one of the following characteristics:
 - (1) substantially 100% of the pores have a pore size less than 0.24 micron;
 - (2) at least 80% of the pores have a pore size less than 0.15 micron;
 - (3) at least 70% of the pores have a pore size less than 0.11 micron;
 - (4) the pores constitute 35 to 80% of the volume of the film;
 - (5) the pores constitute 60 to 75% of the volume of the film:
 - (6) it comprises a polymeric matrix composed of
 - essentially linear ultrahigh molecular weight polyethylene having an intrinsic viscosity of at least 18 deciliters/g, or
 - essentially linear ultrahigh molecular weight polypropylene having an intrinsic viscosity of at least 6 deciliters/g, oi
 - (c) a mixture of (a) and (b);
 - (7) it comprises (i) a polymeric matrix, and (ii) 50 to 90% by weight, based on the weight of the microporous film, of a finely divided, particulate, substantially water-insoluble, siliceous filler which is distributed throughout the matrix; and
 - (8) it has been calendered at a nip pressure of 17.8 to 266.6 kg/linear cm (100 to 1,500 pli) before or after the coating is formed thereon.

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- A membrane according to claim 5, 6 or 7 wherein the coating polymer has at least one of the following characteristics:
 - it is a crystalline polymer having a T_p of -5 to 40°C and a ΔH of at least 5
 J/e:
 - (2) it is a crystalline polymer having a T_p of 0 to 15°C and a ΔH of at least 20 J/g;
 - it is a side chain crystalline polymer;
 - it is a side chain crystalline polymer in which T_p-T_o is less than 10°C;
 - (5) it is a side chain crystalline polymer prepared by copolymerizing (f) at least one n-alkyl acrylate or methacrylate in which the n-alkyl group contains at least 12 carbon atoms and (ii) one or more comnomers selected from acrylic acid, methacrylic acid, and esters of acrylic or methacrylic acid in which the esterifying group contains less than 10 carbon atoms:
- it is cis-polybutadiene, poly(4-methylpentene), polydimethyl siloxane, or ethylene-propylene rubber; and
 - it has been crosslinked.
 - 9. A package which is stored in air and which comprises
 - (a) a sealed container, and
 - (b) within the sealed container, a respiring biological material and a packaging atmosphere around the biological material

the sealed container including one or more permeable control sections which provide substantially the only pathways for oxygen and carbon dioxide to enter or leave the packaging atmosphere, at least one said permeable control section being a gas-permeable membrane as claimed in any one of claims 1 to 4.

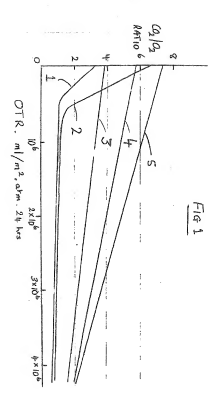
- 10. A package which is stored in air and which comprises
 - (a) a sealed container, and

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 within the sealed container, a respiring biological material and a packaging atmosphere around the biological material

the sealed container including one or more permeable control sections which provide substantially the only pathways for oxygen and carbon dioxide to enter or leave the packaging atmosphere, at least one said permeable control section being a gas-permeable membrane as claimed in any one of claims 5 to 8.

11. A package according to claim 9 or 10 wherein the control section is in the form of a strip which extends across a complete dimension of the container.



6N9DOCID: 4WO 9636495A1 I

INTERNATIONAL SEARCH REPORT

Inte 'onel Application No PCT/US 96/07939

A. CLASSIFICATION OF SUBJECT MATTER IPC 6 C08J9/36 B65D81/24

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation selected (classification system followed by classification symbols) IPC 6 C08J B65D

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim	Relevant to claim No.	
X	US.A.5 254 354 (R.F. STEMART) 19 October 1993 cited in the application see claims 1-16 see examples 1,2,4,7 see tables 3,7,8	1-11		
X	WO,A,94 12040 (FRESH MESTERN MARKETING, INC.) 9 June 1994 see claims 1-6,10 see page 4, line 12-23 see examples 1-3	1-11		
Х	US,A,5 160 768 (M.K.ANTOOM, JR.) 3 November 1992 cited in the application see Claims 1-5 see examples 1-10	1-11		

*Special categories of cited documents: 'A' document defining the general state of the art which is not considered to be of particular relevance.	The face operations patientistical airc the international filting data or principly date and on its position with the application but related to understand the principle or theory industrying the critical to understand the principle or theory industrying the control of the principle or theory industrying the control of principles or the control to deduce the desired to lower the considered to work to extract the observation in the control of the contro			
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Date of the actual completion of the international search 26 September 1996	Date of mailing of the international search report 0 2, 10, 96			
Name and mailing address of the ISA. European Paters Office, P.B. 5818 Patentiam 2 N.I 2280 HV Rijnorit, Tel. (+ 31.70) 340-1040, Tz. 31 651 epo rk, Faze (+ 31.70) 340-3104.	Authorised officer Hallemeesch, A			

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INTERNATIONAL SEARCH REPORT

Int sonal Application No PCT/US 96/07939

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